This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims: Please amend the claims as follows:

We claim:

Claims 1.-53. (Cancelled)

Claim 54. (Currently Amended)

A process for the production of ergosterol or an intermediate product thereof which is squalene, farnesol, geraniol, lanosterol, zymosterol, 4,4-dimethylzymosterol, 4-methylzymosterol, ergost-7-enol or ergosta-5,7-dienol or a sterol with a 5,7-diene structure, comprising fermenting into ergosterol with a plasmid vector-transformed microorganism, wherein said vector comprises several suitable genes of the ergosterol metabolic process in altered form, wherein the catalytic area of HMG is expressed without its membrane-bonded domain; and the natural promoter of t-HMG, ERG9 and SAT1 gene is replaced by the middle part of ADH1 promoter, the suitable genes being

a-i)

- i) a gene of the HMG-Co-A-reductase (t-HMG),
- a gene of the squalene synthetase (ERG9),
- iii) a gene of the acyl-CoA; sterol-acyltransferase (SAT1), and
- iv) a gene of squalene epoxidase (ERG1),

or

a-ii)

- i) a gene of HMG-Co-A-reductase (t-HMG), and
- ii) a gene of the squalene synthetase (ERG9),

or

a-iii)

i) a gene of the HMG-Co-A-reductase (t-HMG), and

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a gene of the acyl-CoA: sterol-acyltransferase (SAT1),
       ii)
or
a-iv)
              a gene of the HMG-Co-A-reductase (t-HMG), and
       i)
       ii)
              a gene of squalene epoxidase (ERG1),
or
a-v)
       i)
              a gene of squalene synthetase (ERG9), and
       ii)
              a gene of acyl-CoA: sterol-acyltransferase (SAT1)
or
a-vi)
       i)
              a gene of squalene synthetase (ERG9), and
       ii)
              a gene of squalene epoxidase (ERG1),
or
a-vii)
       i)
              a gene of acyl-coA: sterol-acyltransferase (SAT1) and
       ii)
             a gene of squalene epoxidase (ERG1),
or
a-viii) at least one of the genes of a-i),
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Claim 55. (Previously Presented) A process according to claim 54, further comprising a gene of squalene epoxidase (ERGI) in a-ii), a-iii), and a-v), and a gene of

analyzing, purifying and isolating the fermented product.

acyl-CoA: sterol-acyl-transferase in a-ii).

Claim 56. (Previously Presented) A process according to claim 54, wherein the genes in a-i) to a-vii) are introduced, in each case independently of one another, into microorganisms of the same species.

and

Claim 57. (Cancelled)

Claim 58. (Cancelled)

Claim 59. (Cancelled)

Claim 60. (Previously Presented) A process according to claim 54, wherein the microorganism is yeast.

Claim 61. (Previously Presented) A process according to claim 60, wherein said yeast is S. errevisiae.

Claim 62. (Previously Presented) A process according to claim 61, wherein said yeast is S. cerevisiae AH22.

Claim 63. (Previously Presented) Yeast strain S. cerevisiae AH22, comprising one or more of the genes mentioned under a-i) of claim 54.

Claim 64. (Cancelled)

Claim 65. (Cancelled)

Claim 66. (Cancelled)

Claim 67. (Cancelled)

Claim 68. (Previously Presented) An expression cassette that comprises the average ADH promoter, the t-HMG gene, the TRP terminator and the SAT1 gene with the average ADH promoter and the TRP terminator.

Claim 69. (Previously Presented) An expression cassette comprising the average ADH promoter, the t-HMG gene, the TRP terminator, the SAT1 gene with the average ADH promoter, and the TRP terminator, and the ERG9 gene with the

average ADH promoter and the TRP terminator.

Claim 70. (Previously Presented) A combination of expression cassettes comprising

 a) a first expression cassette, on which the ADH promoter, the t-HMG gene

and the TRP terminator are located,

a second expression cassette, on which the ADH promoter, the SATI gene

and the TRP terminator are located,

and

 a third expression cassette, on which the ADH promoter, the ERG9 gene
 with the TRP terminator are located.

Claim 71. (Cancelled)

Claim 72. (Cancelled)

Claim 73. (Previously Presented) A microorganism comprising an expression cassette according to claim 68.

Claim 74. (Previously Presented) A microorganism according to claim 73, which is yeast.

Claim 75. (Cancelled)

Claim 76. (Cancelled)

Claim 77. (Previously Presented) A process according to claim 55, wherein the genes are introduced, in each case independently of one another, into microorganisms

of the same species.

Claim 78. (Previously Presented) A microorganism comprising an expression cassette according to claim 69.

Claim 79. (Previously Presented) A microorganism comprising a combination according to claim 70.

Claim 80. (Previously Presented) A microorganism according to claim 78, which is yeast.

Claim 81. (Previously Presented) A microorganism according to claim 79, which is yeast.

Claim 82. (New) A process for the production of ergosterol or an intermediate product thereof which is squalene, farnesol, geraniol, lanosterol, zymosterol, 4,4-dimethylzymosterol, 4-methylzymosterol, ergost-7-enol or ergosta-5,7-dienol or a sterol with a 5,7-diene structure, comprising fermenting into ergosterol with a plasmid vector-transformed yeast, wherein said vector comprises suitable genes of the ergosterol metabolic process in altered form, wherein the catalytic area of HMG-COA reductase (HMG1) is expressed without its membrane-bonded domain; and the natural promoter of altered variant of HMG-1 (t-HMG), squalene synthetase (ERG9) and sterol-acyl transferase (SAT1) gene is replaced by the middle part of ADH1 promoter, the suitable genes being

a-i)

- i) a gene of the HMG-Co-A-reductase (t-HMG),
- ii) a gene of the squalene synthetase (ERG9),
- iii) a gene of the acyl-CoA; sterol-acyltransferase (SAT1), and
- iv) a gene of squalene epoxidase (ERG1),

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or
a-ii)
             a gene of HMG-Co-A-reductase (t-HMG), and
      i)
             a gene of the squalene synthetase (ERG9),
      ii)
or
a-iii)
      i)
             a gene of the HMG-Co-A-reductase (t-HMG), and
      ii)
             a gene of the acvl-CoA: sterol-acvltransferase (SAT1).
or
a-iv)
             a gene of the HMG-Co-A-reductase (t-HMG), and
      i)
      ii)
             a gene of squalene epoxidase (ERG1),
or
a-v)
      i)
             a gene of squalene synthetase (ERG9), and
      ii)
             a gene of acyl-CoA; sterol-acyltransferase (SAT1)
or
a-vi)
      i)
             a gene of squalene synthetase (ERG9), and
      ii)
             a gene of squalene epoxidase (ERG1),
Of
a-vii)
             a gene of acyl-coA: sterol-acyltransferase (SAT1) and
      i)
      ii)
             a gene of squalene epoxidase (ERG1),
or
a-viii) at least one of the genes of a-i),
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analyzing, purifying and isolating the fermented product.

and

Claim 83. (New) The process according to claim 82, wherein said suitable genes of the ergosterol metabolic process are yeast genes.